



# (imiglucerase for injection)

### 400 UNITS

#### DESCRIPTION

is a lysosomal glycoprotein enzyme which catalyzes the hydrolysis of the glycolipid B-Glucocerebrosidase (B-D-glucosyl-N-acylsphingosine glucohydrolase, E.C. 3.2.1.45) Cerezyme® (imiglucerase for injection) is an analogue of the human enzyme glucocerebroside to glucose and ceramide. B-glucocerebrosidase, produced by recombinant DNA technology

specifically recognized by endocytic carbohydrate receptors on macrophages, the cells cerebrosidase. These mannose-terminated oligosaccharide chains of imiglucerase are structures on imiglucerase are somewhat different from those on placental glucosites have been modified to terminate in mannose sugars. The modified carbohydrate histidine is substituted for arginine. The oligosaccharide chains at the glycosylation differs from placental glucocerebrosidase by one amino acid at position 495, where Cerezyme® is produced by recombinant DNA technology using mammalian cell culture that accumulate lipid in Gaucher disease. 497 amino acids, containing 4 N-linked glycosylation sites (Mr = 60,430). Imiglucerase (Chinese hamster ovary). Purified imiglucerase is a monomeric glycoprotein of

Cerezyme® is supplied as a sterile, non-pyrogenic, white to off-white lyophilized product. The quantitative composition of the lyophilized drug is provided in the following table:

| Ingredient   | 200 Unit Vial       | 400 Unit Vial       |
|--|---------------------|---------------------|
| lmiglucerase (total amount)*   | 212 units           | 424 units           |
| Mannitol   | 170 mg              | 340 mg              |
| Sodium Citrates  | 70 mg               | 140 mg              |
| (Trisodium Citrate)<br>(Disodium Hydrogen Citrate)                     | (52 mg)<br>(18 mg)  | (104 mg)<br>(36 mg) |
| Polysorbate 80, NF   | 0.53 mg             | 1.06 mg             |
| Citric Acid and/or Sodium Hydroxide may have been added at the time of | en added at the tim | e of                |
| manufacture to adjust pH.  |                     |                     |

<sup>\*</sup>This provides a respective withdrawal dose of 200 and 400 units of imiglucerase.

1 micromole of the synthetic substrate para-nitrophenyl-ß-D-glucopyranoside (pNP-Glc) per minute at 37°C. The product is stored at 2-8°C (36-46°F). After reconstitution with Sterile Water for Injection, USP, the imiglucerase concentration is 40 U/mL An enzyme unit (U) is defined as the amount of enzyme that catalyzes the hydrolysis of Reconstituted solutions have a pH of approximately 6.1. (see DOSAGE AND ADMINISTRATION for final concentrations and volumes).

# CLINICAL PHARMACOLOGY

# Mechanism of Action/Pharmacodynamics

engorged and are typically found in the liver, spleen, and bone marrow and occasionally resulting in accumulation of glucocerebroside in tissue macrophages which become Gaucher disease is characterized by a deficiency of ß-glucocerebrosidase activity, pathological fractures. Cerezyme® (imiglucerase for injection) skeletal complications, including osteonecrosis and osteopenia with secondary and thrombocytopenia in addition to the characteristic progressive hepatosplenomegaly, in lung, kidney, and intestine. Secondary hematologic sequelae include severe anemia catalyzes

> (alglucerase injection). and decreased cachexia to a degree similar to that observed with Ceredase® the hydrolysis of glucocerebroside to glucose and ceramide. In clinical trials, Cerezyme® improved anemia and thrombocytopenia, reduced spleen and liver size,

### Pharmacokinetics

from placental-derived alglucerase (Ceredase®). level and infusion rate. The pharmacokinetics of Cerezyme® do not appear to be different 3.6 to 10.4 minutes. Plasma clearance ranged from 9.8 to 20.3 mL/min/kg (mean ± S.D., 0.09 to 0.15 L/kg (0.12  $\pm$  0.02 L/kg). These variables do not appear to be influenced by dose or duration of infusion. However, only one or two patients were studied at each dose Following infusion, plasma enzymatic activity declined rapidly with a half-life ranging from During one-hour intravenous infusions of four doses (7.5, 15, 30, 60 U/kg) of Cerezyme® 14.5  $\pm$  4.0 mL/min/kg). The volume of distribution corrected for weight ranged from (imiglucerase for injection), steady-state enzymatic activity was achieved by 30 minutes

elimination half-life compared to patients without antibody (see WARNINGS) In patients who developed IgG antibody to **Cerezyme®**, an apparent effect on serum enzyme levels resulted in diminished volume of distribution and clearance and increased

# INDICATIONS AND USAGE

disease that results in one or more of the following conditions: therapy for pediatric and adult patients with a confirmed diagnosis of Type 1 Gaucher Cerezyme® (imiglucerase for injection) is indicated for long-term enzyme replacement

- thrombocytopenia
- bone disease

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d. hepatomegaly or splenomegaly

## CONTRAINDICATIONS

There are no known contraindications to the use of **Cerezyme®** (imiglucerase for injection). Treatment with **Cerezyme®** should be carefully re-evaluated if there is significant clinical evidence of hypersensitivity to the product.

#### WARNINGS

to Cerezyme® (imiglucerase for injection) during the first year of therapy. Patients who patients with detectable IgG antibodies experienced symptoms of hypersensitivity. developed antibodies to Cerezyme® after 12 months of therapy. Approximately 46% of developed IgG antibody did so largely within 6 months of treatment and rarely Approximately 15% of patients treated and tested to date have developed IgG antibody

Conversely, not all patients with symptoms of hypersensitivity have detectable IgG antibody. It is suggested that patients be monitored periodically for IgG antibody formation during the first year of treatment. Patients with antibody to **Cerezyme®** have a higher risk of hypersensitivity reaction

exhibited symptoms of hypersensitivity to the product. Treatment with Cerezyme® should be approached with caution in patients who have

pretreatment with antihistamines and/or corticosteroids. Further treatment with imiglucerase should be conducted with caution. Most patients Anaphylactoid reaction has been reported in less than 1% of the patient population have successfully continued therapy after a reduction in rate of infusion and

#### PRECAUTIONS

#### General

Cerezyme<sup>®</sup>. No causal relationship with Cerezyme<sup>®</sup> has been established. Patients with respiratory symptoms in the absence of fever should be evaluated for the presence of pulmonary hypertension. Gaucher disease and have been observed both in patients receiving and not receiving injection). Pulmonary hypertension and pneumonia are known complications of have also been observed during treatment with **Cerezyme®** (imiglucerase for In less than 1% of the patient population, pulmonary hypertension and pneumonia

management of patients with Gaucher disease Therapy with Cerezyme® should be directed by physicians knowledgeable in the

Caution may be advisable in administration of Cerezyme® to patients previously Ceredase® or who have exhibited symptoms of hypersensitivity to Ceredase® treated with Ceredase® (alglucerase injection) and who have developed antibody to

# Carcinogenesis, Mutagenesis, Impairment of Fertility

impairment of fertility. effects of Cerezyme® (imiglucerase for injection) on carcinogenesis, mutagenesis, or Studies have not been conducted in either animals or humans to assess the potential

# Teratogenic Effects: Pregnancy Category C

clear and the potential benefit is judged by the physician to substantially justify the risk should not be administered during pregnancy except when the indication and need are administered to a pregnant woman or can affect reproductive capacity. Cerezyme® Animal reproduction studies have not been conducted with Cerezyme® (imiglucerase for injection). It is also not known whether **Cerezyme®** can cause fetal harm when

It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when **Cerezyme**® (imiglucerase for injection) is administered to a nursing woman.

marketing experience. Cerezyme® has been administered to patients younger than 2 Cerezyme® and Ceredase® (alglucerase injection) in adults and pediatric patients, group is supported by evidence from adequate and well-controlled studies of established in patients between 2 and 16 years of age. Use of Cerezyme® in this age years of age, however the safety and effectiveness in patients younger than 2 have not with additional data obtained from the medical literature and from long-term post-The safety and effectiveness of Cerezyme® (imiglucerase for injection) have been

## ADVERSE REACTIONS

the percentages calculated for the frequencies of adverse reactions are most likely Since the approval of Cerezyme® (imiglucerase for injection) in May 1994, Genzyme greater than the actual incidences. over that span of time. The actual number of patients exposed to  ${\sf Cerezyme}^{\otimes}$  since to the voluntary nature of the database and the continuous accrual and loss of patients exposure to **Cerezyme**® since 1994. Actual patient exposure is difficult to obtain due using the number of patients from these sources as the denominator for total patient percentage of events for each reported adverse reaction term has been calculated adverse events and adverse events discussed in the medical literature. has maintained a worldwide post-marketing database of spontaneously reported 1994 is likely to be greater than estimated from these voluntary sources and, therefore,

burning, swelling or sterile abscess at the site of venipuncture. Each of these events was found to occur in < 1% of the total patient population. events were related to the route of administration. These include discomfort, pruritus, administration and which occurred with an increase in frequency. Some of the adverse of patients experienced adverse events which were judged to be related to Cerezyme® Experience in patients treated with Cerezyme® has revealed that approximately 13.8%

Symptoms suggestive of hypersensitivity have been noted in approximately 6.6% of reduced rate of infusion have allowed continued use of Cerezyme® in most patients total patient population. Pre-treatment with antihistamines and/or corticosteroids and reported (see WARNINGS). Each of these events was found to occur in < 1.5% of the dyspnea, coughing, cyanosis, and hypotension. Anaphylactoid reaction has also been these symptoms include pruritus, flushing, urticaria, angioedema, chest discomfort, patients. Onset of such symptoms has occurred during or shortly after infusions;

treated with **Cerezyme®** include: nausea, abdominal pain, vomiting, diarrhea, rash, fatigue, headache, fever, dizziness, chills, backache, and tachycardia. Each of these events was found to occur in < 1.5% of the total patient population. Additional adverse reactions that have been reported in approximately 6.5% of patients

adverse events in children (defined as ages 2-12 years) included dyspnea, fever, nausea, flushing, vomiting, and coughing, whereas in adolescents (>12-16 years) and in adults (>16 years) the most commonly reported events included headache. in the post-marketing database. From this database, the most commonly reported Incidence rates cannot be calculated from the spontaneously reported adverse events

Cerezyme®, transient peripheral edema has been reported for this therapeutic class of In addition to the adverse reactions that have been observed in patients treated with

#### OVERDOSE

Experience with doses up to 240 U/kg every 2 weeks have been reported. At that dose there have been no reports of obvious toxicity

# DOSAGE AND ADMINISTRATION

routine comprehensive evaluations of the patient's clinical manifestations. increase or decrease, based on achievement of therapeutic goals as assessed by dictate that treatment be initiated at a relatively high dose or relatively frequent 2 weeks is the dosage for which the most data are available. Disease severity may 2.5 U/kg of body weight 3 times a week to 60 U/kg once every 2 weeks. 60 U/kg every administration. Dosage adjustments should be made on an individual basis and may Cerezyme® (imiglucerase for injection) is administered by intravenous infusion over 1-2 hours. Dosage should be individualized to each patient. Initial dosages range from

solution may be filtered through an in-line low protein-binding 0.2  $\mu m$  filter during used. DO NOT USE Cerezyme® after the expiration date on the vial. administration. Any vials exhibiting opaque particles or discoloration should not be be inspected visually before use. Because this is a protein solution, slight flocculation Cerezyme® should be stored at 2-8°C (36-46°F). After reconstitution, Cerezyme® should (described as thin translucent fibers) occurs occasionally after dilution. The diluted

On the day of use, after the correct amount of Cerezyme® to be administered to the Sterile Water for Injection, USP. The final concentrations and administration volumes are patient has been determined, the appropriate number of vials are each reconstituted with

|                                       | 200 Unit Vial | 400 Unit Viai |
|---------------------------------------|---------------|---------------|
| Sterile water for reconstitution      | 5.1 mL        | 10.2 mL       |
| Final volume of reconstituted product | 5.3 mL        | 10.6 mL       |
| Concentration after reconstitution    | 40 U/mL       | 40 U/mL       |
| Withdrawal volume                     | 5.0 mL        | 10.0 mL       |
| Units of enzyme within final volume   | 200 units     | 400 units     |
|                                       |               |               |

each vial. The appropriate amount of **Cerezyme®** for each patient is diluted with 0.9% Sodium Chloride Injection, USP, to a final volume of 100 - 200 mL. **Cerezyme®** is shown to be stable for up to 24 hours when stored at 2-8°C. **Cerezyme®**, after reconstitution, has been shown to be stable for up to 12 hours when stored at room temperature (25°C) and at 2-8°C. **Cerezyme®**, when diluted, has been after reconstitution, vials should be promptly diluted and not stored for subsequent use administered by intravenous infusion over 1-2 hours. Aseptic techniques should be used when diluting the dose. Since **Cerezyme®** does not contain any preservative, A nominal 5.0 mL for the 200 unit vial (10.0 mL for the 400 unit vial) is withdrawn from

or decreased to utilize fully each vial as long as the monthly administered dosage bottles. Thus, the dosage administered in individual infusions may be slightly increased small dosage adjustments to be made occasionally to avoid discarding partially used Relatively low toxicity, combined with the extended time course of response, allows remains substantially unaltered.

### HOW SUPPLIED

Cerezyme® (imiglucerase for injection) is supplied as a sterile, non-pyrogenic, lyophilized product. It is available as follows:

Store at 2-8°C (36-46°F).

400 Units per Vial NDC 58468-4663-1 200 Units per Vial NDC 58468-1983-1

U.S. Patent Numbers: 5,236,838 5,549,892

Genzyme Corporation Cerezyme® (imiglucerase for injection) is manufactured by:

500 Kendall Street Cambridge, MA 02142 USA

Certain manufacturing operations may have been performed by other firms

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